

**REQUEST FOR REVIEW OF RESEARCH ACTIVITY INVOLVING HUMAN
SUBJECTS**

INSTRUCTIONS: Please type directly on this form. You can expand the document if you need more space. If your research involves a survey or questionnaire, please attach it to this completed form.

Completed forms (with all required signatures) may be sent to OHSR by FAX (301-402-3443) or by mail (101C116). If you have any questions, call OHSR at (301) 3444.

Date: _____

To: OFFICE OF HUMAN SUBJECTS RESEARCH, Building 10, Room 1C-116

From: _____

Through: _____
(Signature of appropriate Official for IC, e.g., Lab/Branch Chief)

Name of NIH Research Investigator(s): _____

IC _____ Laboratory/Branch _____
Building & Room No. _____ Tel. No. _____ FAX No. _____

Is the NIH research investigator an NIH employee? ____ Yes ____ No

If no, please explain: _____

1. What is the proposed research activity that you intend to perform at NIH (please use lay terms): _____

2. Who is/are your Associate or Collaborating Investigator(s)?

Name	Institution	Address	Tel. #	FAX #
------	-------------	---------	--------	-------

_____	_____	_____	_____	_____
_____	_____	_____	_____	_____
_____	_____	_____	_____	_____
_____	_____	_____	_____	_____

3. Proposed start date of your research:_____Proposed completion date:

4. What role will you have in this research project? (Check all that apply)

- ☐ Analyze samples/data only
☐ Consultant/advisor to collaborator(s) listed above
☐ Author of the protocol that is being implemented by your collaborating investigator (identified in question #2)
☐ Co-authorship on publication(s)/manuscript(s) pertaining to this research
☐ You/NIH hold an IND used in this research

Will you have a decisional authority over the design or implementation of the research at the IRB approved site? Yes_____ No_____

Explanation_____

_____Other (If necessary, use this space to describe your role in this research

5. Where are the subjects of this research activity located?

6. If human subjects are located elsewhere (not at NIH), will you have direct contact or intervention with them? (Examples: as subject's physician; in obtaining samples directly from the subject; by interviewing the subject?) ☐ Yes_____No

7. What kind of human samples (e.g., tissue, blood) or data (e.g., private information, responses to questionnaires) will be involved in your research?

8. Will you be _____these samples or data?

Collecting Yes/No

Receiving Yes/No

Sending Yes/No

9. Do the samples or data:

(a) Already exist? ☐ Yes ☐ No

(b) Or are they being collected for the express purpose of this study?

☐ Yes ☐ No

If "yes," please describe:_____

(c) Or a combination of (a) and (b)? ☐ Yes ☐ No

10. If the samples, data do not come from an IRB approved protocol, do they come from:

- (a) Repository ____ Yes ____ No
- (b) Pathological waste ____ Yes ____ No
- (c) Autopsy material ____ Yes ____ No
- (d) Publicly available source ____ Yes ____ No
- (e) Other _____

11. Will the samples/data that you receive be coded? _____ Yes _____ No

(a) Will your collaborator(s) be able to identify individual subjects? ____ Yes ____ No

(b) Will any information you receive make it possible for you to identify individual subjects? ____ Yes ____ No

. (c) _____ No, the samples are anonymized/unlinked (The samples/data cannot be linked to individual subjects by anyone, including you and your collaborators at the other site(s).)

(d) _____ Other (If other, please explain.) _____

12. Will you send results back to the provider(s) (listed in question 2 of this form)

- (a) ____ No, I will not send results back to the provider(s)
- (b) ____ Yes, I will send aggregate results to the provider(s)
- (c) ____ Yes, I will send coded results to the provider(s)

If yes, does the provider intend to link your data to identifiable individuals?
____ Yes ____ No

13. Has the research activity that you are proposing in this form been approved by an Institutional Review Board(IRB) elsewhere?

____ Yes, the NIH research activity has been reviewed by the following IRB (s)
(Please provide the following information for **each** IRB):

____ Name of institution that provided the review

____ Address of reviewing institution

_____ Name of PI for the IRB approved protocol

_____ Title of IRB approved protocol and protocol #

_____ Federal Wide Assurance (FWA) number**

_____ No IRB review of the research activity described in question #1 above has taken place

(**An FWA is a contract between the U.S. Department of Health and Human Services (DHHS) and an entity receiving DHHS funds to conduct clinical research that the latter will follow ethical guidelines and federal regulations for the protection of human subjects. For a list of domestic and international institutions go to <http://ohrp.cit.nih.gov/search/asearch.asp#ASUR>

14. Per NIH guidance***, have conflicts of interest by NIH employees, if any, been resolved?
_____ Yes _____ No

If your answer is no, please see your Clinical Director about this matter before proceeding with this research.

***The January 5, 2005 NIH Guide to Preventing Conflict of Interest applies to all research conducted at NIH, http://ohsr.od.nih.gov/New/mpafwa_docs.html

6/2/05